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(71) Applicant (<i>for all designated States except GB</i>): FORD MOTOR COMPANY [US/US]; Parklane Towers East, Suite 911, One Parklane Boulevard, Dearborn, MI 48126 (US).		
(71) Applicant (<i>for GB only</i>): FORD MOTOR COMPANY LIMITED [GB/GB]; Eagle Way, Brentwood, Essex CM13 3BW (GB).		
(72) Inventors: JONES, James, Victor; 5255 Snowden Drive, Toledo, OH 43623 (US). BOULOS, Edward, Nashed; 4395 Stonehenge Court, Troy, MI 48098 (US).		
(74) Agent: MESSULAM, Alec, Moses; A. Messulam & Co., 24 Broadway, Leigh on Sea, Essex SS9 1BN (GB).		

(54) Title: A NITRATE-FREE METHOD FOR MANUFACTURING A BLUE GLASS COMPOSITION**(57) Abstract**

The invention is nitrate-free method to manufacture a blue soda-lime-silica glass composition and improving the ultraviolet light absorption thereof while maintaining a high visible light transmission. The method comprises the steps of: adding colorants, during molten glass formation, consisting essentially of a manganese compound along with a cobalt compound, iron oxide and optionally titanium oxide to a soda-lime-silica base glass composition, and adding no sodium nitrate into the batch during molten glass formation, the materials are added in quantities sufficient to form said blue glass composition having at a 4.0 mm. thickness: 477 – 494 dominant wavelength and 6 – 40 % purity of excitation. The blue glass composition, by weight, comprises a base glass composition of: 68 to 75 % SiO₂, 10 to 18 % Na₂O, 5 to 15 % CaO, 0 to 10 % MgO, 0 to 5 % Al₂O₃, and 0 to 5 % K₂O, where CaO + MgO is 6 to 15 % and Na₂O + K₂O is 10 to 20 %; and colorants consisting essentially of: 0.4 to 2.0 % total iron oxide as Fe₂O₃, 0.15 to 2.00 % manganese oxide as MnO₂; 0.005 to 0.025 % cobalt oxide as Co, and 0 to 1.00 % titanium oxide as TiO₂. The glass is useful for automotive or architectural applications. The method preferably involves the use of a reducing agent like anthracite coal during melting operations.

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**A NITRATE-FREE METHOD FOR MANUFACTURING
A BLUE GLASS COMPOSITION**

The invention is directed to a nitrate-free method of
5 manufacturing a blue glass composition having improved
ultra-violet (UV) and infra-red (IR) absorption together
with a low shading coefficient. More particularly, the blue
glass is a soda-lime-silica glass whose colorants are iron
10 oxide, cobalt oxide, manganese oxide and optionally titanium
oxide. The method may use, e.g., anthracite coal as a
reducing agent.

Reference is made to related US patent application
Serial No. 08/891684 entitled "A BLUE GLASS COMPOSITION WITH
15 IMPROVED UV AND IR ABSORPTION" concurrently filed and
commonly assigned herewith.

Blue glass has found particular utility for
architectural applications as building glass and has been
considered for automotive glass applications. Blue glass
has been manufactured using iron oxide, cobalt, and selenium
20 as colorants, the cobalt imparting a blue colour to the
glass, as disclosed in US Patent RE 34,639 to Boulos et al.
In US Patent RE 34,760, Boulos et al disclose blue glass
using iron oxide, cobalt, and nickel, and yet in another
embodiment, a blue glass further including selenium, both
25 patents being commonly owned with the present invention.
Selenium is however a costly colorant and tends to
volatilise from the glass while nickel is prone to
undesirable nickel sulphide "stone" formation within the
glass. Still another blue glass composition is disclosed in
30 US Patent 5,344,798, the glass including iron oxide and
cerium oxide, together optionally with limited amounts of
titanium oxide, zinc oxide, manganese oxide, and cobalt
oxide, and having a specific Fe^{+2}/Fe^{+3} wt. ratio. The use of
cerium oxide, which is known to improve UV absorption, is
35 less than commercially desirable because cerium oxide is
extremely expensive.

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Iron oxide exists in two forms in glass, the reduced (Fe+2) form and the oxidised (Fe+3) form. Another blue glass composition, disclosed in US Patent 3,779,733, incorporates tin oxides to form more of the blue reduced (Fe+2) iron 5 which has IR absorbing properties. This decreases the amount of the yellow oxidised iron, however, which has UV absorbing properties. Others have used a combination of zinc and tin oxides to similarly reduce the iron and form a pale blue glass, as in US Patent 5,013,487, with a resultant lowering 10 of the UV absorption. Another pale blue glass is obtained in US Patent 4,792,536 using a unique melting furnace to reduce the iron oxide, without using tin or zinc oxides, which again lowers the UV absorption.

As would be appreciated, the UV and IR light absorption 15 properties of iron oxide are especially valuable when the glass is to be used in buildings. When heat is absorbed by the glass, the load on building air conditioners is reduced and when the ultra violet absorption is improved, there is less damage over time to the colours of articles inside the 20 building, additionally providing for more comfort.

Therefore, controlling these spectral properties of the glass is very important. Adding iron oxide to the glass under normal furnace conditions improves both the ultra-violet and the infrared absorption of the glass since the 25 concentration of both iron forms is correspondingly increased, but this improvement is at the expense of visible transmittance since the reduced form is darker in colour. That is, as iron oxide is added, the colour of the glass darkens so that the visible transmittance is correspondingly 30 decreased which may limit the usefulness of the glass.

It is unexpectedly found that the present invention blue glass can include relatively large amounts of iron oxide yet have good visible transmittance and excellent UV absorbing properties, without the undesirable aspects of 35 some prior art blue glasses, through the use of a particular combination of colorants in the glass, i.e., iron oxide, cobalt oxide and manganese oxide. For example, the present

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blue glass provides this excellent UV absorption without the use of costly UV absorbers like cerium oxide. In US patent applications Serial No. 08/762,474 filed December 9, 1996 and CIP application Serial No. 08/767,768 filed December 17, 5 1996, both commonly assigned herewith, the present inventors disclosed a high transmittance green glass with improved UV absorption, whose colorants are iron oxide and manganese oxide, and optionally any of titanium oxide, cerium oxide, vanadium oxide, and chromium oxide.

10 The present invention is a nitrate-free method for manufacturing a blue glass composition and improving the ultraviolet light absorption thereof while maintaining a high visible light transmission, which method comprises the steps of: adding colorants, during molten glass formation, 15 consisting essentially of a manganese compound along with a cobalt compound, iron oxide and optionally titanium oxide to a soda-lime-silica base glass composition, and adding no sodium nitrate into the batch during molten glass formation, to manufacture a blue glass. According to the method the 20 materials are added in quantities sufficient to form a blue glass composition whose base glass composition comprises: 68 to 75% SiO_2 , 10 to 18% Na_2O , 5 to 15% CaO , 0 to 10% MgO , 0 to 5% Al_2O_3 , and 0 to 5% K_2O , where $\text{CaO} + \text{MgO}$ is 6 to 15% and $\text{Na}_2\text{O} + \text{K}_2\text{O}$ is 10 to 20%; and whose colorants consist 25 essentially of: 0.4 to 2.0% total iron oxide as Fe_2O_3 ; 0.15 to 2.00% manganese oxide as MnO_2 ; 0.005 to 0.025% cobalt oxide as Co , and 0 to 1.00% titanium oxide as TiO_2 , all by weight percent based on the total weight of the blue glass composition; the glass having at a 4.0 mm. thickness: 477 - 30 494 dominant wavelength and 6 - 40% purity of excitation.

Advantageously, embodiments of the glass composition have improved UV and IR absorption and a low shading coefficient as compared to some other blue glasses of similar colour and visible transmittance. The shading 35 coefficient is a widely used indicator of solar heat gain through a glass when compared to a standard reference glazing system. It is a significant advantage to improve UV

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and IR absorption, in a commercially desirable way without the use of expensive absorbers like cerium oxide as done herein, while maintaining good visible transmittance. It is found that the manganese oxide colorant of the present blue 5 glass composition allows particularly for the avoidance of often added colorants like selenium or nickel oxide, while obtaining a pleasing medium to dark blue colour.

Thus in this method of manufacturing the glass, no sodium nitrate is added during glass melt processing.

10 Anthracite coal, a reducing agent, is optionally but desirably added to the batch materials to aid in refining during the manufacturing process, in particular to cooperate with a conventional fining agent, sodium sulphate. This co-operation lowers the temperature at which SO_3 is released 15 from the sulphate. The anthracite coal may be replaced in part or in total by other reductants like blast furnace slag, slag from coal fired furnaces, coke, or graphite in the batch. In the present invention, the use of such 20 reducing agents can act to override some of the oxidising effect of the manganese oxide colorant or sodium sulphate fining agent without destroying the blue colour of the glass. These and other advantages of the present invention will become apparent from the detailed description which 25 follows.

25 Soda-lime-silica glass, used in the automotive and architectural industries and conveniently made by the float glass process, is generally characterised by the following basic composition shown in Table I, the amounts of the components being based on a weight percentage of the total 30 glass composition:

TABLE I

Base Glass Components	Weight %
SiO ₂	68 to 75
Al ₂ O ₃	0 to 5
CaO	5 to 15
MgO	0 to 10
Na ₂ O	10 to 18
K ₂ O	0 to 5

The blue glass composition made according to the method of the present invention method employs this basic soda-lime-silica glass composition wherein, additionally, CaO + MgO is 6 to 15% and Na₂O + K₂O is 10 to 20%. In addition, the colouring components of the blue glass composition consist essentially of: 0.4 to 2.0 wt. % total iron oxide as Fe₂O₃, 10 0.15 to 2.00 wt.% manganese oxide as MnO₂; 0.005 to 0.025 wt. % cobalt oxide as Co, and 0 to 1.00 wt.% titanium oxide as TiO₂. Further, the invention blue glass considered at a 4.0 mm. thickness has the following spectral properties: 477 to 494 dominant wavelength and 6 to 40% purity of excitation.

Blue glass products made according to embodiments of the invention preferably have the following spectral properties at 4.0 mm. thickness: 20 to 70 % light transmittance using Illuminant A (LTA) and less than 62% ultra violet (UV) transmittance measured over the range of 20 300 to 400 nanometers and less than 54% infra red (IR) transmittance measured over the range of 760 to 2120 nanometers. A preferred embodiment of glasses of the invention include a LTA of less than 60% with the UV transmittance being less than 50% and the IR transmittance 25 being less than 45%. The most preferred embodiment of glasses of the invention include a LTA of less than 50% with the UV transmittance less than 40% and the IR transmittance less than 30%.

Generally, as quantities of the colorants increase in 30 the glass composition, the %LTA, %IR and %UV transmittance

of the glass will go down. Similarly, as the glass thickness increases for a given glass composition, the transmittance of the thicker glass decreases. Preferably, the dominant wavelength is between 480 and 488 nanometers 5 with an excitation purity of 10 to 30%. Most preferably, the dominant wavelength is between 482 and 485 nanometers with an excitation purity of 10 to 20%.

Melting and refining aids are routinely included in glass manufacture and may also be used herein. One refining 10 aid generally used to remove bubbles from the glass is sodium sulphate which generates SO₃ in the glass. Preferably SO₃ is present in the glass composition in a range of 0.10 to 0.30 wt.%, more preferably this range is 0.14 to 0.25 wt.%.

One required colorant of the invention blue glass is 15 iron oxide, wherein as total iron oxide (as Fe₂O₃) it is present in quantities of 0.4 to 2.0 weight %, more preferably being 0.6 to 1.2 weight percent. All weight percents herein being based on the total weight of the invention blue glass composition. Typically, this colorant 20 is added into the batch ingredients in the oxide form, Fe₂O₃. As discussed above, iron oxide exists in two forms in the glass melt. The oxidised form of iron oxide absorbs ultra violet (UV) light and the reduced form of iron oxide absorbs infra red (IR) light, hence the presence of iron oxide 25 lowers the UV and IR transmittance through the glass products. Both absorbing functions of the iron oxide are especially valuable when the glass product is used in architectural applications particularly in geographic areas having a significant amount of sunshine.

30 Another essential colorant in the invention blue glass composition is manganese oxide which is present in the composition in an amount of 0.15 to 2.0 wt.% as MnO₂, more preferably in an amount of 0.2 to 0.8 wt % MnO₂. The manganese component can be added to the batch glass 35 materials in a variety of manganese compound forms, including, but not limited to, MnO, MnO₂, Mn₃O₄, MnSO₄, MnCO₃, MnCl₂, MnF₂, etc, as well as mixtures of any of them. This

colorant is generally present as a Mn ⁺² and Mn ⁺³ oxide of manganese in the glass, although it may additionally or alternately be present in other states such as Mn ⁺⁴. It is generally expected that any manganese compound used would be 5 present in the glass as manganese oxide.

It is important that the Mn₂O₃ form of manganese oxide generally absorbs in the same spectral area as selenium or nickel oxide colorants. Thus it is found it may be used in the blue invention composition to provide, in part, the 10 colouring effect of selenium or nickel oxide in obtaining the desired blue colour of the present invention glass, yet without the drawbacks of either selenium or nickel. As disclosed above, selenium is expensive and easily volatilised from the glass melt. Manganese oxide, for 15 example, is inexpensive and not subject to such volatility as is selenium so that it is optimal as a colorant in the present blue glass composition. Using nickel oxide as a colorant brings about the undesirable potential to form nickel sulphide stones in the glass when sulphates are used 20 as fining agents. Nickel sulphide stones are small ellipsoids that escape normal inspection methods during glass manufacturing and have been known to cause spontaneous breakage upon tempering of the glass. It has often been suggested in the literature that the use of manganese oxide 25 along with iron oxide should be avoided in glass compositions because of the tendency of the glass to then solarise. That is, manganese oxide has been known to cause glass to discolour when it is exposed to a strong ultraviolet light. USP 5,344,798 mentioned above discusses 30 the solarisation problem as related to the inclusion of manganese oxide in the glass and limits its inclusion. In the present invention, the composition includes relatively large amounts of manganese oxide and yet it is found, as shown in the examples, that this amount of manganese oxide 35 is not expected to cause the glass to solarise.

The manganese colorant has oxidising ability which it is found additionally useful in the present invention. We

wish to oxidise the iron oxide towards its more colourless form. Oxidising environments have been provided to glass melts in several ways, for example by providing additional air to the glass melt in the furnace, increasing sodium 5 sulphate, calcium sulphate, or sodium nitrate in the batch or lowering the temperature of the furnace. All of these efforts have commercial drawbacks. For example, sodium nitrate can lead to undesirable nitrogen oxide emissions. It is found that the use of the manganese oxide colorant in 10 the glass in a range of 0.15 to 2.00 wt.% as MnO_2 additionally provides oxidising benefits in the manufacture of the present invention blue glass eliminating the need of any additional oxidiser, like sodium nitrate. Thus, desirably, this blue glass composition may be manufactured 15 without the use of sodium nitrate, as is preferred herein.

When the manganese compound is added to the glass batch, it is reduced to its more colourless form. For example, a portion of the purple coloured manganese oxide colorant in its oxidised form (e.g., Mn_2O_3) is converted to 20 the more colourless, reduced MnO . It is thus found more iron oxide may be added to the batch to enhance both the ultraviolet and the infra red absorption of the glass while simultaneously maintaining a high visible transmittance and obtaining a desired blue colour of the glass. While it is 25 expected that the other manganese compounds like $MnCl_2$ would be similarly useful and converted to oxides in the batch, preferably it is most desirable to use the manganese oxide or manganese carbonate compounds as sources of the manganese oxide colorant in the glass batch.

30 Cobalt is another colorant required in the blue glass composition made according to the present invention method. It is typically added to the batch ingredients as an oxide compound thereof, and is present as a colouring component in the glass in an amount of 0.005 to 0.025 wt. % as Co , 35 preferably in an amount of 0.005 to 0.015 wt. % and most preferably in an amount of 0.006 to 0.012 wt.% as Co . The cobalt colorant functions to absorb light in the 580 to 680

nanometer range of the visible spectrum. The strong absorption in the 580 to 680 nanometer range and weaker absorption in the lower wavelengths is what primarily gives glass of the present invention its blue colour. It is

5 necessary to balance the amount of absorption from MnO_2 and both FeO and Fe_2O_3 with that of cobalt to achieve the desired blue appearance of the present glass composition.

Ordinarily, in conventional glass compositions, increasing the amount of iron oxide would undesirably reduce 10 the amount of visible light being transmitted through the glass. Thus while the UV and IR properties might be improved in conventional glass by increasing the iron oxide colorant, if a high visible light transmittance glass were desired, it would not be achieved. The present invention 15 advantageously provides a blue glass with good UV and IR absorption, while at the same time maintaining good visible transmittance and a pleasing medium blue colour. And it provides the good UV absorption without the use of costly UV absorbers like cerium oxide.

20 One property of the manganese colorant, its oxidising ability toward the iron oxide, acts to improve the UV absorption of the invention blue glass. And, in the present invention, increasing the total iron concentration can improve the IR absorption plus again lower the UV 25 absorption. In another preferred embodiment of the present invention discussed in more detail below, anthracite coal or other reductant is used together in the batch with the colorants to further increase the UV and IR absorption of the glass product. The anthracite coal shifts some of the 30 manganese oxide and iron oxide toward their reduced forms and the effect is to enhance the blue colour. The visible transmittance (%LTA) is lowered when manganese oxide, cobalt oxide, iron oxide and anthracite coal are used in the manner described above. Some of the cobalt oxide can be removed to 35 raise the %LTA and the glass will remain blue from the colouring effect of the reduced species of both iron oxide and manganese oxide.

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The particular amounts of each of the colorant employed, e.g., iron oxide and each form of iron oxide (Fe^{+3} , Fe^{+2}) in a embodiment of a glass product according to the present invention will be a matter of choice and dependent 5 in part on the desired spectral properties of the blue glass product, as would be apparent to those skilled in the art in view of the present disclosure. Selection of a particular embodiment glass composition will be dependent in large part on its desired application, such that one application glass 10 product advantageously has more UV absorption while another glass product optimally has better IR absorption.

As explained above, the method of the present invention produces a glass with improved UV absorption while allowing maintenance of good visible transmittance without the use of 15 costly additives. For example, a sample of a commercially available blue glass made according to US Patent RE 34,639, discussed above and being commonly assigned herewith, includes iron oxide, cobalt, and selenium as colorants and has a UV transmittance of about 64.7 % and an IR 20 transmittance of 48.8 % at 64.7 % LTA (composition of Example 1 herein). A present invention embodiment of similar colour appearance can be made having, at 63.6 % LTA, a UV transmittance of 40.0 % and an IR transmittance of 40.6 % as in Example 3. Example 2 has a UV transmittance of 45.5% and 25 an IR transmittance of 41.0% at 64.7% LTA. These examples demonstrate the significant improvement of the UV and IR properties of embodiments of the present blue composition as compared to available blue glass with almost the same % LTA. The advantage for architectural applications of the blue 30 glass made according to the present invention method is apparent from the comparisons of the spectral properties made above.

The present blue glass composition may also include 35 titanium oxide as TiO_2 in an amount up to 1.0 wt. % to improve the UV absorption of the glass. Generally, the present invention glass does not require any added titanium oxide since it possesses excellent UV and IR properties.

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Should it be desired to enhance the UV absorption, titanium dioxide may be added and preferably when included it will comprise up to about 0.4 wt. % of the blue glass composition.

5 It is known that tramp materials may enter the glass batch during production changeover from one glass composition to another in the glass melting furnaces or from impurities often accompanying the raw materials used. Exemplary of such tramp materials are selenium, nickel
10 oxide, molybdenum, zinc, zirconium, lithium, and chromium, although this list is not meant to be limiting. Still others will be apparent to those skilled in the art in view of the present disclosure. These tramp materials or impurities are expected to be in small amounts, e.g., up to
15 0.0005 wt.% selenium and up to 0.005 wt. % nickel oxide as NiO. Depending on the source of the raw materials, of course, titanium dioxide often enters soda-lime-silica glass compositions as an impurity with the sand, dolomite or limestone in levels that give rise in the final glass
20 product, even when no titanium dioxide has intentionally been added, in a range of about 0.015wt.% or 0.02 wt.% to about 0.05 wt.% titanium dioxide.

25 The following table lists ingredients which are preferably used to form the optimal embodiments of blue glass compositions made according to the present invention.

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TABLE II

BATCH MATERIALS	RANGE MASS (LBS.)
SAND	1000
SODA ASH	290 TO 350
DOLOMITE	215 TO 260
LIMESTONE	70 TO 90
SALT CAKE	6 TO 24
ROUGE (97% Fe ₂ O ₃)	6 TO 30
MANGANESE DIOXIDE	2.0 TO 28
COBALT OXIDE (Co ₃ O ₄)	0.095 TO 0.5
TITANIUM DIOXIDE	0 TO 14
CARBOCITE	0 TO 4
NEPHELINE SYENITE	0 TO 150

In order to demonstrate the advantages of blue glass
5 made according to the present invention, glass melts
detailed in all of the examples were made in the laboratory
according to the following procedure: batches were weighed,
placed into a glass jar about 2" high and 2" inside diameter
and dry mixed for 10 minutes each on a Turbula mixer, dry
10 batch was placed into an 80% platinum/20% rhodium crucible
that stands 2" tall and has an inside diameter at the top of
2.5" and is tapered to the base which has an inside diameter
of 1.75". An amount of 4.5 ml. of water is added to the dry
batch in the crucible and mixed with a metal spoon. After
15 such preparation, a group of six different batches is melted
in a gas/air fired furnace at the same time for 1 hour at
2600°F and each crucible is removed in turn from the furnace
and fritted. Fritting the glass involves coating the inside
of the platinum/rhodium crucible with the molten glass and
20 then plunging the crucible into cold water.

After removing the crucible from the water and draining
the water, the broken glass particles are removed from the
sides of the crucible and mechanically mixed inside the
crucible. All six samples are fritted in like manner and
25 all crucibles are placed back into the furnace for another 1

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hour interval at 2600°F and the fritting procedure is repeated. After the second fritting process, the crucibles are returned to the furnace for 4 hours at 2600°F. Each crucible is removed in turn from the furnace and each molten 5 glass sample is poured into a graphite mould with an inside diameter of 2.5". Each glass is cooled slowly, labelled, and placed into an annealing furnace where the temperature is quickly raised to 1050°F, held for 2 hours, and then slowly cooled by shutting off the furnace and removing the 10 samples after 14 or more hours. The samples are ground and polished to about 4.0 mm. thickness and subsequently the spectral properties are measured for each sample.

All the laboratory melts of the examples are made with the above procedure and use a base composition of 100 grams 15 sand, 32.22 grams of soda ash, 8.81 grams of limestone, 23.09 grams of dolomite, 1.2 grams of sodium sulphate, 2.64 grams of nepheline syenite, and the remainder of the batch includes rouge, manganese dioxide and cobalt oxide and may include anthracite coal or other reductant as described in 20 the example melts. Titanium dioxide also may be added to improve ultra violet absorption, if desired. The base composition of a typical glass melt made from the batch materials above would be about 72 wt.% SiO₂, 13.5 wt.% Na₂O, 0.15 wt.% K₂O, 8.4 wt.% CaO, 3.6 wt.% MgO, 0.6 wt.% Al₂O₃, 25 and 0.2 wt.% SO₃. The range of colorants of the invention embodiment examples herein was: 0.4 to 2.0 wt.% Fe₂O₃, 0.15% to 2.00 wt.% MnO₂, 0.005 to 0.025 wt.% CoO, and 0 to 1.0 wt.% TiO₂, the specific amounts being detailed in the examples. As would be appreciated by those skilled in the art, the wt. 30 % concentration of the base components would decrease as the total amount of colorants is increased.

Table III shows the improvement of the ultra violet and infra red absorption of several examples of embodiments of present invention glass compositions (i.e., other than 35 Example 1 which is a comparative example) including varying amounts of manganese dioxide colorant. In particular, Table III below shows the improvements in ultra violet absorption

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with an increasing amount of MnO_2 at a constant level of 0.6 wt.% Fe_2O_3 . For comparison, Example 1 is a commercially made product based on US Patent RE. 34,639 commonly owned herewith as discussed above, which contains about 0.0002 wt.% selenium. In both Tables III and IV, no TiO_2 was added to the glass, but it was present as an impurity in the glass at a level of about 0.02 wt.%, having come in with raw materials.

10

TABLE III

	Ex. 1	Ex. 2	Ex. 3	Ex. 4	Ex. 5	Ex. 6	Ex. 7
Wt% Fe_2O_3	0.42	0.6	0.6	0.6	0.6	0.6	0.6
Wt% FeO	0.096	0.123	0.125	0.109	0.123	0.254	0.266
ppm Co	50	50	50	150	150	150	150
Wt% MnO_2	None	0.2	0.6	0.2	0.6	0.2	0.6
% LTA	64.7	64.7	63.6	49.3	42.9	40.3	39.3
% UV	64.7	45.5	40.0	42.9	39.3	50.9	46.9
% IR	48.8	41.0	40.6	44.1	40.5	18.2	16.9
% TSET	58.7	53.6	52.3	49.8	45.4	33.1	31.6
Dom. Wave-length	481.9	484.3	486.3	479.6	479.4	478.9	479.4
% Excitation Purity	8.7	9.4	8.2	20.2	23.7	29.8	29.5

From Table III, it can be readily seen that the addition of manganese dioxide colorant together with the relatively increased amount of iron oxide colorant significantly improves both the ultra violet and infra red absorption of the present invention blue glass composition. Particularly, compare the spectral properties of commercial product of Example 1 to glass of the present invention embodiments in Examples 2 and 3. Most desirably, in addition to the significant improvement in the ultra violet absorption of the glass, the invention examples also maintain the visible transmittance of the glass, as evidenced by a similar %LTA. Examples 4 and 5 shows the increase in blue colour intensity by increasing the

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concentration of cobalt oxide with correspondingly significant improvements in both UV and IR absorption. Examples 6 and 7 are similar to Examples 5 and 6 respectively, except that anthracite coal was added to the 5 batch such that the salt cake to anthracite coal ratio was 7:1 to generate reducing conditions. The reducing conditions significantly improved the IR absorption while also attaining better UV absorption than the commercial blue glass product of Example 1.

10 While Table III showed the improvement in ultra violet absorption of present invention glasses with constant total iron colorant when the MnO_2 colorant was increased, Table IV shows the change in ultra violet absorption when a constant amount of MnO_2 (0.2 wt.%) is added to various concentrations 15 of Fe_2O_3 .

The results of Table IV, for present invention blue glass composition embodiments, demonstrate that, in glasses with a constant MnO_2 wt.% (concentration), increasing the Fe_2O_3 , correspondingly increases the ultra violet absorption. 20 Table IV also shows that at a given concentration of MnO_2 colorant, the dominant wavelength (colour) tends to increase slightly as the total iron oxide is increased in the glass. Example 3 (Table III) is the same as Example 9 in Table IV. Example 8 demonstrates the improved UV absorption compared 25 to Example 1 in Table III based, it is believed, on the action of the manganese colorant to shift the iron oxide colorant toward its oxidised form. The blue colour and intensity of present invention embodiment blue glass in Example 8 is similar to the commercial blue glass of Example 30 1 and yet has the added benefit of a higher % LTA. Table IV also demonstrates that the infra red transmittance is lowered in the present invention embodiments as the total iron oxide is increased in the blue invention glass compositions.

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TABLE IV

	Ex. 8	Ex. 9	Ex. 10	Ex. 11	Ex. 12	Ex. 13
Wt.% Fe ₂ O ₃	0.4	0.6	0.75	0.9	1.2	1.6
Wt%FeO	0.091	0.125	0.228	0.318	0.450	0.644
ppm Co	50	50	50	50	50	50
Wt%MnO ₂	0.2	0.2	0.2	0.2	0.2	0.2
% LTA	68.0	64.7	58.9	54.0	46.6	35.1
% UV	56.1	45.5	41.4	36.9	27.7	17.5
% IR	50.4	41.0	21.5	12.7	6.1	1.6
% TSET	60.5	53.6	40.3	33.0	25.4	16.9
Dom. Wave-length	482.7	484.3	485.2	485.8	487.0	488.5
%Excitation Purity	9.4	9.4	12.4	14.4	16.2	19.5

Table V demonstrates the spectral property changes that occur when sodium sulphate and anthracite coal are varied within the embodiment of the invention. The total iron oxide, ppm Co and wt.% MnO₂ were maintained at constant values and the % FeO is the only variable. The actual %Fe₂O₃ varies proportionately with the respective concentration of %FeO to maintain the total iron oxide at 0.6 wt.%. Note that the anthracite coal has a stronger impact on the concentration of %FeO than does the concentration of sodium sulphate. Examples 14 through 17 have the same wt ratio of sodium sulphate to anthracite coal (7:1), but the reducing action of the coal overwhelms the oxidising effect of increasing the sodium sulphate. Note also in Example 16, 18, and 19 at constant sodium sulphate concentration that as the anthracite coal is lowered, then the % FeO gradually falls. In Examples 14 through 19, the constant cobalt oxide and manganese dioxide keep the dominant wavelength and excitation purity within a small range.

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TABLE V

	Ex. 14	Ex. 15	Ex. 16	Ex. 17	Ex. 18	Ex. 19
Wt.% Fe ₂ O ₃	0.6	0.6	0.6	0.6	0.6	0.6
Wt%FeO	0.274	0.303	0.331	0.353	0.282	0.274
ppm Co	80	80	80	80	80	80
Wt%MnO ₂	0.4	0.4	0.4	0.4	0.4	0.4
Na ₂ SO ₄ /1000 SiO ₂	7.5	10	12	16	12	12
Coal/ 1000 SiO ₂	1.07	1.43	1.71	2.29	0.8	0.48
% LTA	51.6	52.3	50.4	48.1	51.6	52.3
% UV	48.5	50.4	51.0	47.7	46.2	50.3
% IR	16.4	13.9	11.7	10.5	15.6	16.3
% TSET	35.4	34.1	32.1	30.0	34.5	35.7
Dom. Wave- length	482.6	483.1	483.1	484.0	483.7	482.5
%Excita- tion Purity	19.3	18.7	19.9	19.1	17.3	19.4

Table VI below further demonstrates the impact of using anthracite coal as compared to not using coal in the batch. Manganese dioxide colorant, sodium sulphate and total iron oxide colorant were kept constant while the % FeO varied due to the reducing power of the coal. Examples 20 through 22 did not have any reducing agent in the batch while Examples 23 through 25 each had anthracite coal in the batch which reduced the iron oxide to generate a higher % FeO and the % IR transmitted was much lower. Note that in each case as the cobalt oxide increased, the dominant wavelength dropped and the % excitation purity increased indicating that the blue colour was more intense.

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TABLE VI

	Ex. 20	Ex. 21	Ex. 22	Ex. 23	Ex. 24	Ex. 25
Wt.% Fe ₂ O ₃	0.6	0.6	0.6	0.6	0.6	0.6
Wt%FeO	0.121	0.115	0.117	0.150	0.148	0.152
ppm Co	65	80	95	65	80	95
Wt%MnO ₂	0.2	0.2	0.2	0.2	0.2	0.2
Na ₂ SO ₄ /1000 SiO ₂	12	12	12	12	12	12
Coal/1000 SiO ₂	0	0	0	0.8	0.8	0.8
% LTA	57.9	57.6	54.2	59.7	56.3	52.6
% UV	41.3	44.9	44.8	46.2	46.5	45.9
% IR	41.2	43.1	42.4	34.6	34.9	34.0
% TSET	51.1	52.3	50.8	48.4	47.5	45.7
Dom. Wave- length	482.8	481.8	480.9	482.9	481.9	481.0
%Excitation Purity	12.9	14.0	16.3	12.9	15.3	18.1

Table VII below demonstrates the increase in the dominant wavelength as the manganese oxide colorant is increased from 0.2 to 0.6 wt.% plus the oxidising effect that lowers the % UV transmitted at the higher manganese oxide concentration and at constant iron oxide levels. It further demonstrates the influence of increased iron oxide colorant. In particular, as the iron oxide concentration increases; the %UV, %IR, and the %LTA are all lowered. All of the Examples in Table VII demonstrate the increased absorption in the IR from the addition of anthracite coal.

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TABLE VII

	Ex. 26	Ex. 27	Ex. 28	Ex. 29	Ex. 30	Ex. 31
Wt.% Fe ₂ O ₃	0.4	0.4	0.9	0.9	1.6	1.6
Wt%FeO	0.163	0.160	0.373	0.367	0.622	0.644
ppm Co	150	150	150	150	150	150
Wt%MnO ₂	0.2	0.6	0.2	0.6	0.2	0.6
Na ₂ SO ₄ /1000 SiO ₂	12	12	12	12	12	12
Coal/ 1000 SiO ₂	1.71	1.71	1.71	1.71	1.71	1.71
% LTA	43.8	43.2	36.4	36.1	24.6	23.3
% UV	61.6	55.2	37.8	35.6	16.4	12.7
% IR	31.5	32.1	9.1	9.4	2.0	1.9
% TSET	42.5	42.1	25.2	25.0	13.9	12.9
Dom. Wave- length	477.9	478.4	480.4	480.9	482.8	483.9
%Excita- tion Purity	28.5	27.3	30.1	29.4	31.7	30.1

Blue glass compositions made according to the present invention can be used for both automotive and architectural applications. Generally, they would be manufactured by well known float glass techniques. Present federal automotive regulations require a minimum of 70.0% LTA measured at the actual glass thickness, in general, for glazings used in automotive vehicles. For example, sun-roofs may have a lower LTA. The blue glass of the present invention is expected to maintain its LTA throughout its useful life. Glasses containing manganese and iron oxide colorants have been known to solarise or discolour when exposed to a strong ultra violet light source as discussed above. Tests of glass compositions in the ranges of the concentrations of the iron oxide with manganese oxide colorants of the present blue glass compositions by the inventors have demonstrated that solarisation does not take place. It is expected that

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the present invention blue glass additionally including cobalt oxide will likewise not experience solarisation since it is the manganese oxide which has been repeatedly implicated in literature as causing solarisation, not cobalt oxide.

5 The examples demonstrate the inventors unexpected discovery of an advantageous way to improve ultra violet light absorption in blue glass products while maintaining good visible transmission and, at the same time, improving 10 absorption in the infra red portion of the spectrum. All of this has remarkably been accomplished in a low cost and environmentally friendly way, i.e. by using, as one of the colorants, a manganese compounds instead of commonly used materials like selenium, nickel, or cerium oxide in the 15 glass composition batch. And preferably, the blue glass can avoid the inclusion of sodium nitrates with the glass batch as is generally used in glass manufacture, since these nitrates are a source of undesirable NOx emissions.

20 While embodiments of the novel method of making the blue glass composition of the invention has been illustrated and described, it will be obvious to those skilled in the art in the light of the present disclosure that various 25 modifications within the scope of the disclosure may be made without departing from the invention. It is intended to cover in the appended claims all such modifications and equivalents as fall within the true spirit and scope of the invention.

CLAIMS

1. A nitrate-free method for manufacturing a blue glass composition and improving the ultraviolet light absorption thereof while maintaining a high visible light transmission, which method comprises the steps of:

5 adding colorants, during molten glass formation, consisting essentially of manganese compound with cobalt compound, iron oxide and optionally titanium oxide to a 10 soda-lime-silica base glass composition, and adding no sodium nitrate into the batch during molten glass formation, all base materials and colorants being used in sufficient quantities to manufacture a blue glass composition whose base glass composition comprises: 68 to 15 75% SiO_2 , 10 to 18% Na_2O , 5 to 15% CaO , 0 to 10% MgO , 0 to 5% Al_2O_3 , and 0 to 5% K_2O , where $\text{CaO} + \text{MgO}$ is 6 to 15% and $\text{Na}_2\text{O} + \text{K}_2\text{O}$ is 10 to 20%; and whose colorants consist 20 essentially of: 0.4 to 2.0% total iron oxide as Fe_2O_3 ; 0.15 to 2.00% manganese oxide as MnO_2 ; 0.005 to 0.025% cobalt oxide as Co_2O_3 , and 0 to 1.00% titanium oxide as TiO_2 , all by weight percent based on the total weight of the blue glass composition; the glass having at a 4.0 mm. thickness: 477 - 494 dominant wavelength and 6 - 40% purity of excitation.

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2. A method according to claim 1, wherein the dominant wavelength at 4.0 mm. thickness is between 480 and 488 nanometers.

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3. A method according to either claim 1 or claim 2, wherein the purity of excitation at 4.0 mm. thickness is between 10 and 30%.

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4. A method according to any one of the preceding claims, wherein the amount of manganese colorant in the blue glass composition as MnO_2 is 0.2 to 0.8 wt.%.

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5. A method according to any one of the preceding claims, wherein the amount of Fe_2O_3 in the blue glass composition is in the range of 0.6 to 1.2 wt.%.

5 6. a method according to any one of the preceding claims, wherein the amount of cobalt oxide in the blue glass composition as Co is in the range of 0.005 to 0.015 wt.%.

10 7. A method according to any one of the preceding claims, wherein it further comprises floating molten glass made according to the method on a molten tin bath.

15 8. A method according to any one of the preceding claims, which further comprises using during the melt processing a reductant as a raw material component.

20 9. A method according to claim 8, wherein said reductant is selected from the group consisting of anthracite coal, blast furnace slag, slag from coal fired furnaces, coke, or graphite, or mixtures thereof.

10. A method according to claim 8, wherein said reductant comprises at least anthracite coal.

25 11. A method according to any one of the preceding claims, wherein said blue glass composition comprises SO_3 in an amount of between 0.10 and 0.30 wt. %.

30 12. A method according to claim 3, wherein a purity of excitation of the blue glass is from 10 to 20%.

35 13. A method according to any one of the preceding claims, wherein the amount of cobalt oxide as Co in said blue glass composition is within the range of 0.006 to 0.012 wt. %.

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14. A method according to any one of the preceding claims, wherein the dominant wavelength of said blue glass composition is between 482 and 485 nanometers.

5 15. A method according to any one of the preceding claims, wherein said blue glass composition has the following spectral properties at 4.0mm: 20 to 70% light transmittance using Illuminant A (LTA) and less than 62% ultraviolet (UV) transmittance measured over the range of 10 300 to 400 nanometers and less than 54% infra red (IR) transmittance measured over the range of 760 to 2120 nanometers.

15 16. A method according to claim 1, wherein said blue glass composition comprises: 68 to 75% SiO_2 , 10 to 18% Na_2O , 5 to 15% CaO , 0 to 10% MgO , 0 to 5% Al_2O_3 , and 0 to 5% K_2O , where $\text{CaO} + \text{MgO}$ is 6 to 15% and $\text{Na}_2\text{O} + \text{K}_2\text{O}$ is 10 to 20%; and colorants consisting essentially of: 0.6 to 1.2% total iron oxide as Fe_2O_3 ; 0.2 to 0.8% manganese oxide as MnO_2 ; 0.005 to 20 0.015% cobalt oxide as Co_2O_3 , 0 to 1.00% titanium oxide as TiO_2 , all weight percents being based on the total weight of the blue glass composition; the blue glass having at a 4.0 mm. thickness: 480 - 488 dominant wavelength and 10 - 30% purity of excitation.

INTERNATIONAL SEARCH REPORT

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A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 C03C4/02 C03C3/087 C03C4/08

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 6 C03C

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 5 346 867 A (JONES JAMES V ET AL) 13 September 1994 see column 2, line 7 – column 5, line 16 ----	1-16
A	US 5 521 128 A (JONES JAMES V ET AL) 28 May 1996 see column 2, line 8 – column 5, line 11 ----	1-16
A	EP 0 619 274 A (PILKINGTON PLC ;FLACHGLAS AG (DE)) 12 October 1994 see claims ----	1-16
A	WO 97 23422 A (CORNING INC ;BROCHETON YVES A H (FR); COMTE MARIE J M (FR)) 3 July 1997 see claims ----	1-16 -/-

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Name and mailing address of the ISA

European Patent Office, P.B. 5518 Patentlaan 2
NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl.
Fax: (+31-70) 340-3016

Authorized officer

Van Bommel, L

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C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	EP 0 677 492 A (FORD MOTOR CO) 18 October 1995 see page 6 see page 3, line 14 - line 23 -----	1-16
P, A	WO 98 25863 A (FORD MOTOR CO ;FORD MOTOR CO (US)) 18 June 1998 see page 1, line 1 - page 10, line 36 -----	1-16

INTERNATIONAL SEARCH REPORT

Information on patent family members

International Application No

PCT/GB 98/01826

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
US 5346867	A 13-09-1994	CA 2179146 A CN 1137785 A, B DE 69405011 D DE 69405011 T EP 0734358 A ES 2105870 T WO 9516641 A JP 9506584 T US 5521128 A		22-06-1995 11-12-1996 18-09-1997 08-01-1998 02-10-1996 16-10-1997 22-06-1995 30-06-1997 28-05-1996
US 5521128	A 28-05-1996	US 5346867 A CA 2179146 A CN 1137785 A, B DE 69405011 D DE 69405011 T EP 0734358 A ES 2105870 T WO 9516641 A JP 9506584 T		13-09-1994 22-06-1995 11-12-1996 18-09-1997 08-01-1998 02-10-1996 16-10-1997 22-06-1995 30-06-1997
EP 0619274	A 12-10-1994	DE 4311180 C AT 142995 T DE 59400657 D US 5656560 A		12-01-1995 15-10-1996 24-10-1996 12-08-1997
WO 9723422	A 03-07-1997	FR 2742745 A EP 0869925 A		27-06-1997 14-10-1998
EP 0677492	A 18-10-1995	CA 2143968 A JP 7267675 A		11-09-1995 17-10-1995
WO 9825863	A 18-06-1998	US 5776845 A AU 7846498 A		07-07-1998 03-07-1998

INTERNATIONAL SEARCH REPORT

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C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	EP 0 677 492 A (FORD MOTOR CO) 18 October 1995 see page 6 see page 3, line 14 – line 23 -----	1-16
P,A	WO 98 25863 A (FORD MOTOR CO ;FORD MOTOR CO (US)) 18 June 1998 see page 1, line 1 – page 10, line 36 -----	1-16

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Information on patent family members

International Application No

PCT/GB 98/01826

Patent document cited in search report	Publication date	Patent family member(s)			Publication date
US 5346867	A 13-09-1994	CA 2179146 A	CN 1137785 A, B	DE 69405011 D	22-06-1995
		DE 69405011 T	EP 0734358 A	ES 2105870 T	11-12-1996
		EP 0734358 A	ES 2105870 T	WO 9516641 A	18-09-1997
		ES 2105870 T	WO 9516641 A	JP 9506584 T	08-01-1998
		JP 9506584 T	US 5521128 A	US 5521128 A	02-10-1996
					16-10-1997
					22-06-1995
					30-06-1997
					28-05-1996
US 5521128	A 28-05-1996	US 5346867 A	CA 2179146 A	CN 1137785 A, B	DE 69405011 D
		DE 69405011 T	EP 0734358 A	ES 2105870 T	22-06-1995
		EP 0734358 A	ES 2105870 T	WO 9516641 A	11-12-1996
		ES 2105870 T	WO 9516641 A	JP 9506584 T	18-09-1997
		JP 9506584 T			08-01-1998
					02-10-1996
					16-10-1997
					22-06-1995
					30-06-1997
EP 0619274	A 12-10-1994	DE 4311180 C	AT 142995 T	DE 59400657 D	US 5656560 A
		DE 59400657 D	US 5656560 A		12-01-1995
		US 5656560 A			15-10-1996
					24-10-1996
					12-08-1997
WO 9723422	A 03-07-1997	FR 2742745 A	EP 0869925 A		
		EP 0869925 A			27-06-1997
					14-10-1998
EP 0677492	A 18-10-1995	CA 2143968 A	JP 7267675 A		
		JP 7267675 A			11-09-1995
					17-10-1995
WO 9825863	A 18-06-1998	US 5776845 A	AU 7846498 A		
		AU 7846498 A			07-07-1998
					03-07-1998